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(54) 【発明の名称】 排気ガス浄化用触媒

(57)【要約】

(37) 【契例】 (日的) ペロプスカイト型複合酸化物を触媒成分とし、アルミナと共存させることによって高温でも高い比素面積を維持できるようにするととは、LaAIOの全成を抑えて触媒作用を維持させるようにする。 【構成】 ペロプスカイト型構造の複合酸化物に、ZrOを抵加したアルミナと、Pdなどの貴金属とを共存させる。さらに、Ce及びZrを少なくとも含む耐熱性酸化物を共存させてもよい。

【特許請求の範囲】

【請求項1】 一般式Ln; × A×MO;(LnはCeを除く希土類金属、AはCe又はアルカリ土類金属、M は遷移金属で、いずれも1種又は2種以上、O<×<1)で示されるペロプスカイト型構造の複合酸化物と、乙rO。を添加したアルミナをよれませたまで大排気が多くとも共存させた排気ガス浄化用触媒。

【請求項2】 Ce及び2r、又はさちにCe以外の希 土類金属を含む、少なくとも一部が複合酸化物又は固溶 体となっている割熱性酸化物をさらに共存させた請求項 10に記載の排気ガス浄化用触媒。

【発明の詳細な説明】

[0001]

【産業上の利用分野】本発明は800℃より高温で用いても一酸化炭素(CO)、炭化水素(HC)及び酸化窒素(NOx)の浄化能力に優れた排気ガス浄化用三元糖域に関するものである。

[0002]

発明が解決しようとする課題】ペロブスカイト型複合 酸化物をアルミナと共存させた場合、自動車排気ガス中 や大気中で800℃程度の高温で使用していると、ペロ プスカイト構造の組成中のLaがアルミナと反応してL aA10を生成することが判明した。このLaA10,50

は触媒活性を特たず、これが生成することによって触媒 作用の優れたペロブスカイト型結晶構造が破壊される間 題が生じる。そこで、本発明はペロブスカイト型複合酸 化物を触媒の分とし、アルミナと共存させることによっ て高温でも高い比表面積を維持できるようにするととも に、LaA1Oの生成を抑えて触媒作用を維持させる ようにした触媒を提供することを自的とするものであ る。

[0005]

【課題を解決するための手段】本発明の触媒は、一般式 Lncx AxMO, (LntiCeを除く希土類金属、A はCe又はアルカリ土類金属、Mは遷移金属で、いずれ も1種又は2種以上、0くx×1)で示されるペロプス ルイト型構造の複合酸化物と、ZrO。を添加したアル ミナと、貴金属とを少なくとも共存させた排気ガス浄化 用触媒である。この触媒にはCe及びZr、又はさらに Ce以外の希土類金属を含む、少なくとも一部が複合酸 化物又は固溶体となっている耐熱性酸化物をさらに共存 させることができる。この耐熱性酸化物を300でを越 える高温において焼結を抑制し、高い比表面積を維持す る作用をする。

[0006] 本発明の触媒は、例えば、2r〇を添加 したアルミナと一般式Ln-xAxMOパで示されるペ ロブスカイト型構造の複合酸化物とを少なくとも含む複 合酸化物又は固溶体に、食金属を担持させて製造するこ とができる。その担特方法は、pHを4以下又は10よ リ大幸く関製した貴金属塩木溶液を用いて貴金属を含浸 又は吸着させて担持させ、軟燥後に焼成する方法であ る。

○・ 【0007】本発明では、アルミナとペロブスカイト型 複合酸化物とを共存させて複合酸化物又は固溶体とする 前に、アルミナに Z г の。を添加してアルミナ表面に Z r O。は1800でを越える高温においてアルミナ表面の Z r O。は1800でを越える高温においてアルミナの物質 移動を抑制し、ペロブスカイト型核合酸化物中の La と の反応による La A 1 O。少生成を抑制する。これによ り、他媒作用の優れたペロブスカイト型結晶構造は A 1 の」との共存下において、800でを越える高温状態に おいても結晶構造破壊されることなく、触媒活性を維 持することができる。

【0008】アルミナや副熱性酸化物は一般に触媒成分の担体として用いられているのと同様の状態 (形状、粒度、純度、比表面積) で用いればよい。例えば、比表面積は触媒成分を高分散状態に保持するため、20m½。 展以上が望ましい。耐熱性酸化物中のCe、Zr、希土類金属の比率は特に制限されないが、Ceの100原子に対してZrは5~100原子、好ましくは5~50原子、Ce以外の希土類金属は0~100原子、好ましくは5~30原子の原子比となるように構成する。

50 【0009】アルミナや耐熱性酸化物による効果は、触

媒成分をアルミナや耐熱性酸化物上に担持して用いる場 合、及び触媒成分粉末とアルミナや耐熱性酸化物粉末と を混合して使用する場合のいずれの場合にも有効に発揮 され、800℃を越える高温で長時間使用した後でも触 媒成分は担体上に高分散状態に維持されているか、又は アルミナや耐熱性酸化物粉末間に高分散の状態で存在し ている。

【0010】ペロプスカイト型複合酸化物それ自体も三 元触媒として使用できるが、NOxに対する浄化能力が HCやCOに対する浄化能力に比べてやや劣っている。 そこで、NOxに対する活性を高めるためにパラジウム などの貴金属を少量添加した。貴金属はプロプスカイト 型複合酸化物上及びアルミナ上に、又はさらに耐熱性酸 化物上にも担持されている。担持された貴金属は反応す るペロプスカイト型複合酸化物上では固溶及び析出現象 が起こり、一方、反応しない高比表面積のアルミナや耐 熱性酸化物上では析出状態で高分散に担持されることに より、排気ガス変動雰囲気下における800℃を越える 高温でも貴金属の焼結が起きにくく、浄化活性が高度に 維持される。

【0011】本発明における触媒成分の1つであるペロ プスカイト型複合酸化物の割合は触媒全量の1~80重 量%が望ましい。1重量%より少ない場合は触媒活性が 低く、逆に80重量%より多くなるとアルミナや耐熱性 酸化物の効果が現われにくくなり好ましくない。ペロブ スカイト型複合酸化物の形状、粒度、純度、比表面積な どは触媒成分として通常用いられる状態であればよい。 【0012】他の触媒成分である貴金属は白金族のR u、Rh、Pd、Os、Ir及びPtのうちから選ばれ た1種又は2種以上を用いる。これらの貴金属のうちP 30 d を用いた場合にNOx浄化特性が最も向上する。 貴金 属の量は触媒全量に対し0.01~5重量%、望ましく は0.1~2重量%がよい。貴金属が0.01重量%よ り少ない場合はNOx浄化能力の向上が不十分であり、 逆に5重量%を越えてもNOx浄化能力が飽和する。こ れら貴金属は従来の三元触媒において通常用いられてい る状態 (形状、粒度、純度、比表面積) で用いる。この ように、本発明にかかる触媒は、800℃を越える高温 でも耐久性のある安価な排気ガス浄化用三元触媒とな

【0013】本発明の触媒を製造する方法で貴金属を共 存させる工程では、貴金属塩水溶液のpHを4以下とす る製造方法の場合は、水溶性貴金属塩としてはPdCl i、PtCli、RuCli 3HiOなどの塩化物、Pd (NOs)s、Ru(NOs)s、Rh(NOs)sなどの硝酸塩、 Pd(NO1)1(NH1)1、Pt(NO1)1(NH1)1などのジ ニトロジアミン塩など、水溶液が強酸性を示すものが好

【0014】貴金属塩水溶液のpHを10より大きくし て製造する方法の場合は、テトラアミンパラジウムジク 50 ペロブスカイト型複合酸化物(LasiCesi)(Fess

ロライドPd(NH,),Cl:やテトラアミンバラジウム 水酸塩 P d (NH,) (OH) などの塩基性水溶液にアン モニア水や酸を添加してpH>10になるように調製し て用いるか、PdCls、PtCls、RuCls3Hs Oなどの塩化物、Pd(NOs)s、Ru(NOs)s、Rh (NO),などの硝酸塩、又はPd(NO,)(NH,),、P t (NO.).(NH.).などのジニトロジアミン塩などの酸 性水溶液にアンモニア水を添加してpH>10になるよ うに調製して用いる。本発明の触媒は、これにパインダ 一を添加し、所定の形状に成形して用いたり、又は水を 加えてスラリー状として基材に塗布して用いることもで きる。

[0015]

【発明の効果】本発明ではペロブスカイト型構造の複合 酸化物に、ZrOiを添加したアルミナと、貴金属とを 少なくとも共存させたので、800℃を越える高温でも ベロブスカイト型結晶構造が破壊されることなく、活性 を維持することができる。

[0016] 【実施例】

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(実施例1)

手順A: アルミナの前処理

市販のγ-A1:0:粉末100重量部にイオン交換水5 0 重量部を加え、撹拌して含浸させる。次に、硝酸ジル コニル (ZrO:分40.12wt%) 49. 9重量部を イオン交換水50重量部に溶解し、先に含水させた y -A 1:O:粉末に加えて撹拌し、4 0℃で3 0分間保持し て吸着させる。その後、110℃に昇温させたオープン 中で乾燥させる。30分ごとによく撹拌し、水分を飛散 させた後、110℃で12時間乾燥させる。乾燥した粉 末を、電気炉を用いて大気中で600℃で3時間焼成し た後、乳鉢で180 µ m以下の大きさに粉砕し、微細な ZrOiを表面に分散させたAliOi粉末を得た。Ali O₃とZrO₂の重量比は10:2であった。

【0017】<u>手順B</u> : 耐熱性酸化物の製造 アルミナとともに担体として用いることのある耐熱性酸 化物は、市販の高比表面積の酸化セリウム粉末(CeO :比表面積130 m²/g、純度99.9%/TREO (全希土類酸化物)) 111.9gを用意し、これにオ キシ硝酸ジルコニウム(ZrO(NO;);)水溶液(液比重 1.51、液中にZrO:換算で25.0重量%含まれ る) 147.9g、及び硝酸イットリウム(Y(NO_i)_i) 水溶液(液比重1. 62、液中にY:O:換算で21. 7 重量%含まれる) 26.0gを加え、よく撹拌して混合 しながら110℃で10時間大気中で乾燥した。その 後、大気中で600℃で3時間焼成を行ない、(Cea «Z г » » Y ». ю) О:複合酸化物を約150 g 得た。 【0018】<u>手順C</u> : ペロブスカイト型複合酸化物

結晶粉末の製造方法

Cos.) 〇.粉末の調製方法を設明する。硝酸ランタン 103.9 g、硝酸サリウム26.1 g、硝酸コバルト3 4.9 g、硝酸酸 12.7 g を地水に溶解した水溶液の、3リットルを用意した。次に、中和共沈剤として炭酸ナトリウム50 g を溶解した水溶液の、5リットルを用意した。中和共沈剤を先の水溶液に滴下し、共沈物を得した。その共沈物を十分水充し、濾過した後、真空乾燥した。これを600℃で3時間大気中で焼成後、粉砕し、その後、800℃で3時間大気中で焼成後不分からに粉砕し、(LasiCes)(FesiCos)の粉末を作製した。

【0019】<u>手順D</u> : ペロブスカイト型複合酸化物の相特

手順Cで製造したペロプスカイト型複合酸化物粉末75 重量部、手順Aで得た2 F Q 処理済みのA I c の 砂末 6 0 重量部、手順B で製造した耐熱セリア粉末15 重量 8 たりアゾル (固形分10 w t %) 5 0 重量部 (固形 分では5 重量部) 及びジルコニアゾル (固形分3 0 w t %) 3 .3 重量部 (固形分では1 重量部) を、全固形 分が5 0 w t %となるようにイオン水を5 8.7 重量部 加え、ボールミルにより1 2 時間混合しながら粉砕して スラリーを得た。このスラリーをコージェライトハニカ ムに流入させた後、余剣のスラリーを吹き払い、均一に コーティングした。スラリーコート後のかニカムを1 2 0 でで1 2 時間乾燥し、空気中で0 0 でで3 時間焼成 担てエカム状サンブルを得た。焼液後のサンブルの地 特量はハニカムなサンブルを得た。焼液後のサンブルのよ 特量はハニカムなサンブルを得た。焼液後のサンブルのよ

【0020】 手順臣: 貴金属Pdの担持 硝酸パラジウム溶液(Pd激度4.4 w %)をPd分 で1.67重量部となるように38重量部計量し、イオ ン交換水50重量部を加え、pHc2(実測値はpH= 1.8)に調製した。このパラジウム溶液に手順口で得*

* たハニカム状サンブルを浸漬し、40℃で2時間保持してPdを吸着させた。その後、120℃で12時間乾燥させた後、空気中で600℃で3時間焼成し、実施例1の輪峻就料を得た。

【0021】(実施例2) 実施例1における手順Dで、 ペロプスカイト型複合酸化物的末を75重量部とし、2 rO処理済みA1の約末を40重量部とし、耐熱セ リア粉末を35重量部とする他は、実施例1と同様の機 作により実施例2の触媒試料を得た。

○ 【0022】(実施例3)実施例1における手順Dで、ベロブスカイト型複合酸化物粉末を50重量部とし、Z 「〇処理済みA1の粉末を50重量部とし、配熟セリア粉末を50重量部とし、再順EでのPd分を0.8 3重量部となるように18.9重量部の硝酸ペラジウム溶液を計量する他は、実施例1と同様の操作により実施例3の機能軟料を得た。

【0023】(実施例4)実施例1における手順Dで、 ZrO処理済みA1の約末を75重量部とし、耐熱 セリア粉末の添加を省いた他は、実施例1と同様の操作 により実施例4の触媒試料を得た。

【0024】(比較例a) 実施例1の手順Dにおいて、 ZrO処理済みA1の必束を未処理のA1の粉末 に代えた他は実施例1と同様の操作により比較例aの触 健軟料を得た。

【0025】(比較例も)すでに実用化されている自動 車用触媒であるPtーRh/Al:() 触媒を比較例もの 触媒試料とした。PtーRhら者量は0.54重量部で あった。実施例及び比較例の触媒仕様を表1に示し、そ れぞれの触媒活性の測定結果を表2に示す。表2は初期 及び耐入試験後の50%件化温度を示したものである。 [0026]

【表1】

)パラジウム溶液に手順Dで得 *	[XI]		
ペロブスカイト型複合酸化物	アルミナ	耐熱セリア	貴金属
(Lao. sCeo. 2) (Feo. sCoo. 4) 03 [75]	処理すみ(60)	(CeZrY)02(15)	Pd(1.67)
(Lao. eCen. z) (Feg. eCop. 4) Os (50)	処理すみ[50]	(CeZrY)0 ₂ [50]	Pd(0.83
			Pd(1.67
			Pd(1.67
			(0.54
	ペロブスカイト型複合酸化物 (Lao. sCeo. 2) (Feo. sCoo. 4) Og (75) (Lao. sCeo. 2) (Feo. sCoo. 4) Og (75) (Lao. sCeo. 2) (Feo. sCoo. 4) Og (50) (Lao. sCeo. 2) (Feo. sCoo. 4) Og (75)	(Lao. sCeo. 2) (Feo. sCoo. 4) Os(75) 処理すみ(50) (Lao. sCeo. 2) (Feo. sCoo. 4) Os(75) 処理すみ(50) (Lao. sCeo. 2) (Feo. sCoo. 4) Os(75) 処理すみ(50) (Lao. sCeo. 2) (Feo. sCoo. 4) Os(75) 処理すみ(75) (Lao. sCeo. 2) (Feo. sCoo. 4) Os(75) 処理すみ(75) (Lao. sCeo. 2) (Feo. sCoo. 4) Os(75) 処理な人(50)	

[]内の数値は重量部を表わす。

·							
	初期50%浄化温度(℃)			耐久後50%净化湿度(℃)			
Ī	co	нс	NOx	со	нс	NOx	
実施例1	150	162	162	230	233	222	
実施例2	161	180	168	231	242	220	
実施例3	167	171	172	2 4 5	239	247	
実施例4	171	178	180	262	246	253	
比較何a	175	201	197	283	284	268	
比較例b	192	207	197	285	298	281	

【0028】触媒活性の測定と耐久試験は以下のように 行なった。

触媒活性の測定

ハニカム状(セル数400/inch[®])コージェライト担 体 (確径30mm、長き50mm)に担持されたそれぞ れの飲料を下記のモデルガスにて活性を測定した。ガス 温度は機嫌への入口ガス温度で示し、金温から昇温し、* *NO、CO、HC (C,H++C,H) のそれぞれが初期 譲度の50%に低下した温度を50%浄化温度とする。 【0029】また、リッチガスとリーンガスはそれぞれ 1秒毎に切り換えた。 触媒を通るガス流の空間速度 (S V) は30,000/時間とした。 【0030】

	リッチガス	リーンガス
co	2.6 %	0.7 %
HC(C:換算濃度)	0.19%	0.19%
H:	0.87%	0.23%
CO:	8 %	8 %
NO	0.17%	0.17%
O ₁	0.65%	1.8 %
H ₁ O	10 %	10 %
N.	残部	残部

【0031】耐久試験

上記のリッチガスとリーンガスを5秒毎に切り換えて9 30 00℃で30分、750℃で30分のサイクルを15回 繰り返して耐久試験を行なった。耐久試験後にも前記の 方法で能媒活性を測定した。本発明による触媒はいずれ も比較例よりも50%神化複度が低く、触媒活性に優れ ている。また、耐久試験後の触媒のX線回折による測定※

※の結果、比較例 a では触媒成分であるペロブスカイト型) 複合酸化物とアルミナが反応して生じたL a A I O ・() C P D S カードN o. 3 1 - 0 0 2 2) が検出されたが、各実施例の測定結果からはL a A I O は検出されなかった。この結果から、アルミナのZ r O 処理により、高温での劣化反応が抑制されていることが分か。



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(54) CATALYST FOR PURIFICATION OF EXHAUST GAS

PURPOSE: To inhibit the formation of LaAlO3 and maintain catalytic action and maintain a large specific surface area even at high

temp. by allowing a perovskite type multiple oxide as a catalytic component to coexist with alumina. CONSTITUTION: A multiple oxide having a perovskite type structure is allowed to coexist with ZrO2 added alumina and a noble metal such as Pd or to further coexist with a heat resistant oxide contg. at least Ce and Zr.

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JAPANESE [JP,07-068175,A]

CLAIMS DETAILED DESCRIPTION TECHNICAL FIELD PRIOR ART EFFECT OF THE INVENTION TECHNICAL PROBLEM MEANS EXAMPLE

[Translation done.]

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CLAIMS

[Claim(s)]

[Claim 1] The catalyst for exhaust gas purification which made the multiple oxide of the perovskite type structure shown by general formula Ln1-xAxMO3 (the rare earth metal except Ce and A are transition metals, and Ln is [Ce or alkaline earth metal, and M] all one sort or two sorts or more, and 0 < x<1), the alumina which added ZrO2, and noble metals live together at least.

[Claim 2] Ce and Zr, or the catalyst for exhaust gas purification according to claim 1 that made the heat-resistant oxide which contains rare earth metals other than Ce further, and with which at least the part serves as a multiple oxide or the solid solution live together further.

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DETAILED DESCRIPTION

[Detailed Description of the Invention]

[Industrial Application] Even if it uses this invention at an elevated temperature from 800 degrees C, it relates to the three way component catalyst for exhaust gas purification excellent in the purification capacity of a carbon monoxide (CO), a hydrocarbon (HC), and nitrogen oxide (NOx).

[Description of the Prior Art] Utilization is expected as a cheap three way component catalyst for exhaust gas purification with which the multiple oxide which has the perovskite type structure which consists of a rare earth metal, an alkaline earth metal, and transition metals purifies CO, HC, and NOx (refer to JP,59-87046 A and JP,60-82138 A). However, the perovskite mold multiple oxide catalyst aims at using it below 800 degrees C, and when it is necessary to need high catalytic activity and and it also needs to satisfy the endurance in an elevated temperature in a pyrosphere 900 degrees C or more like the catalyst for automobile exhaust, it cannot be said as sufficient catalyst. That is, it will sinter, if a perovskite mold multiple oxide is used at an elevated temperature 900 degrees C or more, and an effective-surface product decreases, and catalytic activity falls remarkably.

[0003] Moreover, although CO and the purification capacity of HC are excellent, the perovskite mold multiple oxide is a little inferior in the purification capacity of NOx, and they are not enough to present practical use as a three way component catalyst for automobile exhaust. Then, in order to improve NOx purification capacity, it is known that what is necessary is just to make noble metals live together in a perovskite mold multiple oxide. On the other hand, the alumina (aluminum oxide; aluminum 203) is known as an outstanding wash coat ingredient which can maintain a high specific surface area also in an elevated temperature 900 degrees C or more, and is widely used by the precious metal catalyst. Then, in order to consider as the three way component catalyst which can be used for a long time by high temperature which calls a perovskite mold multiple oxide catalyst 900 degrees C or more, it is made to coexist with an alumina and it is possible to add noble metals.

[Problem(s) to be Solved by the Invention] When a perovskite mold multiple oxide was made to coexist with an alumina and it was used at the about 800-degree C elevated temperature in motor exhaust and atmospheric air, it became clear that La under presentation of a perovskite structure reacted with an alumina, and generated LaAlO3. This LaAlO3 does not have catalytic activity, but when this generates, the problem by which the perovskite mold crystal structure which was excellent in the catalysis is destroyed produces it. Then, this invention uses a perovskite mold multiple oxide as a catalyst component, and while enabling it to maintain a high specific surface area also at an elevated temperature by making it coexist with an alumina, it aims at offering the catalyst suppress [catalyst] generation of LaAIO3 and it was made to maintain a catalysis.

[0005]

[Means for Solving the Problem] The catalyst of this invention is a catalyst for exhaust gas purification which made the multiple oxide of the perovskite type structure shown by general formula Ln1-xAxMO3 (the rare earth metal except Ce and A are transition metals, and Ln is [Ce or alkaline earth metal, and M] all one sort or two sorts or more, and 0< x<1), the alumina which added ZrO2, and noble metals live together at least. Ce and Zr, or the heat-resistant oxide that contains rare earth metals other than Ce further and with which at least the part serves as a multiple oxide or the solid solution can be made to live together further for this catalyst. This heatresistant oxide also controls sintering in the elevated temperature exceeding 800 degrees C, and carries out the operation which maintains a high specific surface area.

[0006] The catalyst of this invention can make the multiple oxide or the solid solution which contains at least the multiple oxide of the perovskite type structure shown by the alumina and general formula Ln1-xAxMO3 which added ZrO2 able to support noble metals, and can be manufactured. The support approach is the approach of using the noble-metals salt water solution which prepared pH more greatly than 4 or less and 10, making noble metals sink in or adsorb, making support, and calcinating after desiccation. [0007] In this invention, before making an alumina and a perovskite mold multiple oxide live together and considering as a multiple oxide or the solid solution, the description is in the point of adding ZrO2 to an alumina and making an alumina front face distributing ZrO2. ZrO2 on the front face of an alumina controls the mass transfer of an alumina in the elevated temperature exceeding 800 degrees C. and controls generation of LaAlO3 by the reaction with La in a perovskite mold multiple oxide. Thereby, the perovskite mold crystal structure which was excellent in the catalysis can maintain catalytic activity, without destroying the crystal structure also in the

elevated-temperature condition which exceeds 800 degrees C under coexistence with aluminum 203. [0008] What is necessary is just to use an alumina and a heat-resistant oxide in the condition (a configuration, grain size, purity, specific surface area) same with generally being used as support of a catalyst component. For example, more than 20m2/g of specific surface area is desirable in order to hold a catalyst component in the high distribution condition. Although especially the ratio of Ce in a heat-resistant oxide, Zr, and a rare earth metal is not restricted, Zr constitutes five to 50 atom, and rare earth metals other than Ce zero to 100 atom preferably five to 100 atom to 100 atoms of Ge so that it may become the atomic ratio of five to 30 atom.

[0009] When the effectiveness by the alumina or the heat-resistant oxide supports a catalyst component on an alumina or a heatresistant oxide and it uses. And it is demonstrated effectively [in any / in the case of mixing and using catalyst component powder, and an alumina and heat-resistant oxide powder / case]. Also after carrying out long duration use at the elevated temperature exceeding 800 degrees C, the catalyst component is maintained by the high distribution condition on support, or exists in the condition of high distribution between an alumina or heat-resistant oxide powder.

[0010] Although it can be used as a three way component catalyst also in itself [perovskite mold multiple oxide], the purification capacity over NOx is a little inferior compared with the purification capacity over HC or CO. Then, in order to raise the activity over NOx, little addition of the noble metals, such as palladium, was carried out noble metals — a PUROBUSU kite mold multiple oxide and alumina top — or it is further supported also on the heat-resistant oxide. When the supported noble metals are supported with a deposit condition by high distribution on the alumina of high specific surface area which does not react on the other hand, or a heatresistant oxide by dissolution and a deposit phenomenon happening on the perovskite mold multiple oxide which reacts, also at the elevated temperature exceeding 800 degrees C under an exhaust gas fluctuation ambient atmosphere, sintering of noble metals cannot

break out easily and purification activity is maintained by altitude. [0011] I - 80% of the weight of the catalyst whole quantity of the rate of the perovskite mold multiple oxide which is one of the catalyst components in this invention is desirable. If catalytic activity is low and increases more than 80 % of the weight conversely when fewer than 1 % of the weight, the effectiveness of an alumina or a heat-resistant oxide is hard coming to appear and is not desirable. The configuration of a perovskite mold multiple oxide, grain size, purity, specific surface area, etc. should just be in the

condition of usually being used as a catalyst component. [0012] The noble metals which are other catalyst components use one sort chosen from among Ru, Rh, Pd, Os(es), Ir(s), and Pt(s) of a platinum group, or two sorts or more. When Pd is used among these noble metals, an NOx purification property improves most. The amount of noble metals has 0.1 - 2 desirably good % of the weight 0.01 to 5% of the weight to the catalyst whole quantity. Even if improvement in NOx purification capacity is inadequate when there are few noble metals than 0.01 % of the weight, and it exceeds 5 % of the weight conversely, NOx purification capacity is saturated. These noble metals are used in the condition (a configuration, grain size, purity, specific surface area) of usually being used in the conventional three way component catalyst. Thus, the catalyst concerning this invention turns into a cheap three way component catalyst for exhaust gas purification which is durable also at the

elevated temperature exceeding 800 degrees C. [0013] At the process which makes noble metals live together by the approach of manufacturing the catalyst of this invention In the case of the manufacture approach which makes pH of a noble-metals salt water solution four or less As a water-soluble noble-metals salt, chlorides, such as PdCl2, PtCl2, and RuCl3.3H2O, Dinitro diamine salts of that a water solution indicates strong acid nature to be, such as nitrates, such as Pd (NO3)2, Ru (NO3)3, and Rh (NO3)3, Pd(NO2)2(NH3) 2, and Pt(NO2)2(NH3) 2, etc. are desirable. [0014] In the case of the approach of making pH of a noble-metals salt water solution larger than 10, and manufacturing it Tetrapod amine palladium dichloride Pd(NH3)4Cl2 and tetra-amine palladium oxalate (NH3) (OH) Pd 4 [whether it prepares and uses so that aqueous ammonia and an acid may be added in basic water solutions, such as 2, and it may be set to pH>10 and] Nitrates, such as chlorides, such as PdCl2, PtCl2, and RuCl3.3H2O, and Pd (NO3)2, Ru (NO3)3, Rh (NO3)3, Or it prepares and uses so that aqueous ammonia may be added to aqueous acids, such as dinitro diamine salts, such as Pd(NO2)2(NH3) 2 and Pt(NO2)2(NH3) 2, and it may be set to pH>10. The catalyst of this invention adds a binder to this, it can fabricate and use for a predetermined configuration, or it can add water, can make it the shape of a slurry, and can also apply and use it for a base material.

[Effect of the Invention] Activity can be maintained without destroying the perovskite mold crystal structure also at the elevated temperature exceeding 800 degrees C, since the alumina which added ZrO2 to the multiple oxide of perovskite type structure, and noble metals were made to live together at least in this invention.

[0016]

[Example]

Procedure A: The ion-exchange-water 50 weight section is added to the gamma-aluminum 203 powder 100 weight section of pretreatment marketing of an alumina, and is agitated and infiltrated into it. Next, the zirconium nitrate (ZrO 2-minute wt(s)[40.12] %) 49.9 weight section is dissolved in the ion-exchange-water 50 weight section, and it holds for 30 minutes and is made to agitate in addition to 20gamma-aluminum 3 powder which carried out water previously, and to adsorb at 40 degrees C. Then, it is made to dry in the oven which carried out the temperature up to 110 degrees C. After agitating well every 30 minutes and dispersing moisture, it is made to dry at 110 degrees C for 12 hours. After calcinating the dry powder at 600 degrees C in atmospheric air for 3 hours using an electric furnace, the mortar ground in magnitude of 180 micrometers or less, and 20aluminum3 powder which made the front face distribute detailed ZrO2 was obtained. The weight ratio of aluminum2O3 and ZrO2 was 10:2.

[0017] Procedure B: Heat-resistant oxide which has used as support with the manufacture alumina of a heat-resistant oxide, Commercial cerium oxide powder of high specific surface area (2/g CeO2 specific surface area of 130m) 99.9% of purity and TREO(all rare earth oxides)111.9g are prepared. To this, 147.9g (contained 25.0% of the weight by ZrO2 conversion in liquid density 1.51 and liquid) of oxy-zirconium-nitrate (ZrO2 (NO3)) water solutions, And 26.0g (contained 21.7% of the weight by 20Y3 conversion in liquid density 1.62 and liquid) of nitric-acid yttrium (Y(NO3) 3) water solutions was added, and it dried in 10-hour atmospheric air at 110 degrees C, having agitated well and mixing. Then, baking was performed at 600 degrees C in atmospheric air for 3 hours, and about 150g of O(Ce<SUB>0.65Zr 0.30Y0.05)2 multiple oxides was obtained.

[0018] Procedure C: The preparation approach of manufacture approach perovskite mold multiple oxide (La0.8Ce0.2) (Fe0.6Co0.4) O3 powder of perovskite mold multiple oxide crystal powder is explained. 103.9g of lanthanum nitrates, 26.1g of cerium nitrates, 34.9g of cobalt nitrates, and 0.3l. of water solutions which dissolved 72.7g of iron nitrate in pure water were prepared. Next, 0.5l. of water solutions which dissolved 50g of sodium carbonates as a neutralization coprecipitater was prepared. The neutralization coprecipitater was dropped at the previous water solution, and the coprecipitate was obtained. The vacuum drying was carried out, after rinsing the

coprecipitate enough and filtering it. This was ground after baking in 3-hour atmospheric air at 600 degrees C, and it calcinated in 3-hour atmospheric air at 800 degrees C after that, and ground further, and the powder of O(Fe(La0.8Ce0.2)0.6Co0.4) 3 was produced. For [0019] Procedure D.: The perovskite mold multiple oxide powder 75 weight section nanufactured in support procedure C of a perovskite mold multiple oxide, The aluminum203 powder 60 weight section [finishing / ZrO2 processing] obtained in Procedure A, the heat-resistant Seria powder 15 weight section manufactured in Procedure B, They are 58.7 weight ******* and a ball mill about ion the heat-resistant Seria powder 15 weight section manufactured in Procedure B, They are 58.7 weight section (solid content 15 weight section) solids may become 50wt(s)% about the ceria sol (solid content 10wt)% 50 weight section). Solid content 5 weight sections sol (solid content 30wt%) 3.3 weight section (solid content 1 weight section). Mixing for 12 hours, it ground sand the slurry was obtained. The excessive slurry was blown off and homogeneity was coated, after making this slurry flow into a cordierite honeycomb. The honeycomb behind a slurry coat was dried at 120 degrees C for 12 hours, it calcinated at 600 degrees C in air for 3 hours, and the honeycomb-like sample was obtained. The amount of support of the sample after baking was 180g per

noneycome capacity of the support palladium nitrate solution (Pd concentration 4.4wt%) of noble metals [0020] Procedure E: 38 weight sections measuring of the support palladium nitrate solution (Pd concentration 4.4wt%) of noble metals [0020] Procedure of the support of the section was Pd was carried out so that it might become the 1.67 weight section by part for Pd, the ion-exchange-water 50 weight section was pd was interested to pHt 2 (an actual measurement is pH=1.8). It was immersed, the honeycomb-like sample obtained in Procedure added, and it prepared to pHt 2 (an actual measurement is pH=1.8). It was immersed, the honeycomb-like sample of that 120 degrees C D in this palladium solution was held at 40 degrees C for 2 hours, and Pd was made to adsorb. Then example 1 was obtained.

C for 12 hours, it calcinated at 600 degrees C in air for 3 hours, and the catalyst sample of an example 1 was obtained.

C for 12 hours, it calcinated at 600 degrees U in air for 3 hours, and the catalyst sample of an example 1 has been seen and the formal formal (Example 2) in the procedure D in an example 1, made perovskite mold multiple oxide powder into 75 weight sections, made 20processed [ZrO2] aluminum2 powder into 40 weight sections, and the heat-resistant Seria powder was made into 35 weight sections, and also the catalyst sample of an example 2 was obtained by the same actuation as an example 1 in an example 1.

[0022] (Example 3) Perovskite mold multiple oxide powder was made into 50 weight sections in the procedure D in an example 1, 20processed [2rO2] aluminum3 powder was made into 50 weight sections, the heat-resistant Seria powder was made into 50 weight sections, and the palladium nitrate solution of the 18.9 weight sections was measured so that it might become the 0.83 weight section sections, and the palladium nitrate solution of the 18.9 weight sections was measured so that it might become the 0.83 weight sections, and 10023 (Example 4) In the procedure E and also the catalyst sample of an example 3 was obtained by the same actuation as an example 1, made 20processed [2rO2] aluminum3 powder into 75 weight sections, and 10023 (Example 4) In the procedure D in an example 1, made 20processed [2rO2] aluminum3 powder into 75 weight sections, and 10023 (Example 4) In the procedure D in an example 1, made 20processed [2rO2] aluminum3 powder into 75 weight sections, and 10023 (Example 4) In the procedure D in an example 1, made 20processed [2rO2] aluminum3 powder was obtained by the same actuation as an example 1.

[0024] ((a) The example of a comparison) In the procedure D of an example 1, 20processed [Zr02] aluminum3 powder was replaced with 20aluminum3 unsettled powder, and also the catalyst sample of the example a of a comparison was obtained by the same with 20aluminum3 unsettled powder, and also the catalyst sample of the example a of a comparison was obtained by the same

(0025) ((b) The example of a comparison) 20Pt-Rh/aluminum3 catalyst which is a catalyst for automobiles already put in practical use [0025] ((b) The example of a comparison. The Pt-Rh content was the 0.54 weight section. The catalyst was made into the catalyst sample of the example b of a comparison. The Pt-Rh content was the 0.54 weight section. The catalyst specification of an example and the example of a comparison is shown in Table 1, and the measurement result of each catalytic activity is shown in Table 2. Table 2 shows 50% purification temperature after the first stage and a durability test.

[0026]

Table 1]	ペロブスカイト型複合酸化物	アルミナ	耐熱セリア	貴金属
実施例1	(Lao. sCeo. 2) (Feo. 6Coo. 4)03[75]	処理すみ[60]	(CeZrY)O ₂ (15)	Pd(1.67
	(Lao. aCeo. 2) (Feo. 6Coo. 4)O3 (75)			
	(Lao. aCeo. 2) (Feo. 6Coo. 4)O3 (50)			
実施例4	(Lao. sCeo. 2) (Feo. 6Cop. 4) O3 [75]			Pd(1.6
比較例a	(Lao. sCeo. 2) (Feo. 6Cop. 4) 03 (75)			Pd(1.6
比較例b		処理なし[100]		
			1	(0.5

[]内の数値は重量部を表わす。

[0027] [Table 2]

初期50%浄化湿度(℃)			耐久後50%浄化温度(℃)			
co	нс	NOx	со	нс	NOx	
150	162	162	230	233	222	
161	180	168	2 3 1	242	220	
167	171	172	245	239	247	
171	178	180	262	246	253	
175	201	197	283	284	268	
192	207	197	285	298	281	
	CO 150 161 167 171	CO HC 150 162 161 180 167 171 171 178 175 201	CO HC NOx 150 162 162 161 180 168 167 171 172 171 178 180 175 201 197	CO HC NOx CO 150 162 162 230 161 180 168 231 167 171 172 245 171 178 180 262 175 201 197 283	CO HC NOx CO HC 150 162 162 230 233 161 180 168 231 242 167 171 172 245 239 171 178 180 262 246 175 201 197 283 284	



[0028] Measurement and the durability test of catalytic activity were performed as follows.

Activity was measured for each sample supported by the measurement honeycomb-like (number of cels 400-/inch 2) cordierite support (the diameter of 30mm, die length of 50mm) of catalytic activity by the following model gas. The inlet gas temperature to a catalyst shows gas temperature, it carries out a temperature up from a room temperature, and makes purification temperature temperature to which each of NO, CO, and HC (C3H6+C3H8) fell to 50% of initial concentration 50%. [0029] Moreover, rich gas and lean gas were switched for every second, respectively. Space velocity (SV) of the gas stream which

passes along a catalyst was made into 30,000-/time amount.

Rich gas Lean gas CO 2.6 % 0.7 % HC (C1 conversion concentration) 0.19% 0.19% H2 0.87% 0.23% CO2 8 % 8 % NO 0.17% 0.17% O2 0.65% 1.8 % H2O 10% 10 % N2 Remainder Remainder [0031] The rich gas and lean gas of the durability test above were switched every 5 seconds, it repeated at 900 degrees C for 30 minutes, the cycle of 30 minutes was repeated 15 times at 750 degrees C, and the durability test was performed. Catalytic activity was measured by the aforementioned approach also after the durability test. 50%, each catalyst by this invention has low purification temperature, and excels the example of a comparison in catalytic activity. Moreover, although LaAlO3 (JCPDS card No.31-0022) which the perovskite mold multiple oxide and alumina which are a catalyst component reacted, and was produced was detected in the example a of a comparison as a result of measurement by the X diffraction of the catalyst after a durability test, LaAlO3 was not detected from the measurement result of each example. From this result, ZrO2 processing of an alumina shows that the degradation reaction in an elevated temperature is controlled.

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